

We describe here the expression, purification, solid state NMR sample preparation, and initial structural and functional data for three membrane proteins from *Mycobacterium tuberculosis* (Mtb). The three proteins are FtsX, Rv0008c and Rv1861. Solid state NMR is uniquely able to characterize protein structure in a liquid crystalline lipid bilayer environment. We have used N terminal His tag for protein purification. Nickel-NTA chromatography was performed using a semi automated FPLC instrument. Purified ¹⁵N labeled proteins were eluted into 0.2% (Rv0008c and Rv1861) and 0.4% (FtsX) solution of dodecylphosphocholine (DPC) detergent. The approximate protein yield were 45mg/l (Rv0008c), 50mg/l (FtsX) and 25mg/l (Rv1861) respectively. Samples for solid state NMR were prepared by removing the detergent from the purified protein by exhaustive dialysis against 10mM Tris-HCl (pH-8.0) coincident with reconstitution into lipid bilayers. To prepare aligned samples, pelleted liposomes were layered on to thin glass slides and stacked. 30-35 glass slides were hydrated in a process called 'wet stacking' followed by sealing them into a rectangular glass cell. 400 and 600 MHz magnets were used to determine the 1D and 2D spectra of these aligned samples such that the bilayer is parallel to the applied magnetic field direction. FtsX is an ABC transporter containing 4 transmembrane helices (TMH) and its interaction with FtsZ participate in cell division. Rv1861 has 3 TMH and is known to hydrolyze ATP. It forms a stable octameric structure that is presumably facilitated by the GxxxG, GxxxA, and AxxxA sequences in the transmembrane stretches. Rv0008c is a Mtb membrane protein and participates in cell division. It has been found previously in our laboratory that Rv0008c interacts with Rv0011c and this interaction along with other membrane proteins can facilitate the Mtb cell division process.

3370-Pos

First Structural Characterization of a Bon-Domain in a Protein from *Mycobacterium tuberculosis*: OmpATb Tracks toward an Oligomerization Process to form a Cell Wall Pore

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The etiological agent of tuberculosis (TB), *Mycobacterium tuberculosis*, causing nearly two millions deaths per year is presently one of the greatest infectious agents of mortality worldwide. One of the major problems in TB therapy is the slow uptake of drugs across the thick mycobacterial cell wall made of unique lipid and glycolipid moieties. Water-filled protein channels, called porins, are considered as the main pathway for hydrophilic drugs through the mycobacterial cell wall. In this study, we investigated the 3D structure of the porin-like protein OmpATb (326 residues) from *Mycobacterium tuberculosis* by NMR in solution. We have found that the N-terminal domain of OmpATb (73-204), sufficient to form channels in planar lipid bilayers, forms an α/β sandwich composed of 6 β -strands and 3 α -helices. It appears that a sub-domain of this structure is related to the BON (bacterial OsmY and nodulation) domain fold which was initially identified in bacterial proteins as a conserved ~ 60 residues module supposed to associate with phospholipids. Thus our study gives rise for the first time to the 3D fold of a BON domain member. Other bacterial proteins belonging to macromolecular complexes of the type III secretion systems have BON-like domain structures and form multi-subunits membrane-associated rings at the basal body of the secretion machinery. By analogy with these superstructures, we have built a model of an oligomeric ring assembly of the OmpATb protein to support the formation of functional pores in the mycobacterial cell wall. The numbers of units involved in the pore structure is discussed regarding the biochemical and channel properties of OmpATb.

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SP-C Palmitoylation is Crucial for Stabilizing Cholesterol-Containing Surfactant Films during Continuous Compression/Expansion Cycling

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Cholesterol is critical to maintain a dynamic lateral structure in pulmonary surfactant membranes, including a defined fluid-ordered/fluid-disordered phase equilibrium and proper lateral sorting of surfactant proteins and lipids. However, an excess of cholesterol has been linked to impaired surface activity both in surfactant models and in surfactant from injured lungs. Surfactant protein C (SP-C), the smallest and most hydrophobic of all surfactant proteins, has been shown to interact with cholesterol and dual palmitoylation of its N-terminal segment has been shown to drive association with ordered phases in model membranes. Furthermore, it has been proposed that native palmitoylated SP-C

can act in concert with surfactant protein B (SP-B) to permit cholesterol-containing surfactant films to reach very low surface tensions upon compression. In the present work, we report that palmitoylation of SP-C is important for its ability to counteract deleterious effects of cholesterol on surfactant film stability under continuous expansion/compression cycling, as evaluated in a captive bubble surfactometer (CBS) setup. Presence of 5% cholesterol impairs significantly the stability under quasi static and dynamic compression of films composed of DPPC/POPC/POPG/SP-B (50:25:15:1, w/w/w/w), which are able to reach tensions below 3 mN/m with only 20% compression and almost no hysteresis in the absence of cholesterol. Incorporation in the films of 2% native palmitoylated SP-C could alleviate these deleterious effects. However, recombinant non-palmitoylated SP-C was not able to reproduce the stabilizing effect of native SP-C, confirming that palmitoylation of SP-C at its N-terminal end is crucial for its potential function of stabilizing surfactant films during the respiratory cycles in the lung.

3372-Pos

New Insights into the Interfacial Activation of Secreted Phospholipase A2

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Despite numerous studies towards elucidation of the structural basis of activation of secreted PLA2s upon membrane binding (interfacial activation), no consistent or clear picture has emerged thus far. Previously we have reported significant changes in the secondary and dynamic structures of human group IB and IIA PLA2s, as well as changes in their mode of membrane binding during activation. Here we have conducted atomic resolution NMR studies on free and phospholipid micelle-bound human group IIA PLA2 (hIIAPLA2) to detect more detailed molecular events underlying interfacial activation. Two-dimensional ¹H,¹⁵N-HSQC spectra have been obtained at 600 MHz on Ca²⁺-free and Ca²⁺-loaded hIIAPLA2 in the presence of dodecylphosphocholine (DPC) micelles. Upon complex formation with the micelles, signals from arginine side chain -NH₂ groups of Ca²⁺-loaded hIIAPLA2 are observed, whereas for Ca²⁺-free PLA2 these signals are absent because of fast H/D exchange with the solvent. This suggests that the Ca²⁺-loaded hIIAPLA2 tightly binds to the micelles so these groups are sequestered at the PLA2-micellar interface and shielded from the solvent, or that they are otherwise stabilized by strong hydrogen bonding in the micelle-bound state. TROSY experiments (900 MHz) on Ca²⁺-loaded, ¹⁵N,¹³C-labeled hIIAPLA2 in the absence and presence of DPC micelles (1:600 protein-to-DPC molar ratio) identify substantial conformational changes in PLA2 upon binding to the micelles. Based on the assigned chemical shifts, important structural changes occur throughout the protein. The molecular mechanism of the strong increase in activity of hIIAPLA2 upon phospholipid surface binding is likely to involve a widening of the substrate binding pocket, mediated by a rigid-body movement of the N-terminal helix via interactions of the cationic residues (e.g., Arg7) with lipid phosphate groups. This mechanism will be tested in further studies and may be shared by other secreted PLA2 isoforms.

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Peptide-Membrane and Peptide-Peptide Interactions between Myelin Basic Protein and Myelin-Like Lipids Revealed by Covalently Attached Vibrational Labels

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Cyanylation of cysteine reveals the local solvent environment of the modified side chain and the ps-time scale dynamics of membrane-protein interactions via the infrared absorbance of the covalently attached CN probe vibration. Single-cysteine mutants of the primary membrane binding sequence from the myelin basic protein, were synthesized and cyanylated. A combination of transmission and horizontal attenuated total reflectance (HATR) infrared spectra are used to investigate the dynamics of the membrane-peptide interactions. The SCN-labeled side chains report on the geometry of the peptide-membrane binding interface. They also provide information about the reversible aggregation of the peptide, which occurs at high concentration in solution and when in contact with a myelin-like membrane system.

3374-Pos

Structural Characterization of Human Peripheral Myelin Protein 22 Using NMR

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Peripheral myelin protein 22 (PMP22) is a 160-residue integral membrane protein with four putative transmembrane spans. PMP22 is a major protein of peripheral nervous system (PNS) myelin, where its importance is underscored by the fact that heritable mutations in this protein result in Charcot-Marie-Tooth